Table 1

_	_			_
1,00+1,00	דווכץ כת	Polynucleotide ID	2515666CB1	
Doliming 00+1.do	FOT VIIDCITACITAG	SEQ ID NO:	C1	
) + 1,5 ° L	TIICYCE	Polypeptide ID	2515666CD1	
ם בילימים ו	anthadatas	SEQ ID NO:		
Transfer   Dollary	TIICYCE	Project ID	2515666	

Table 2

GenBank Probability GenBank ID NO: Score	g181382 6.90E-270 Cytochrome P450 [Homo sapiens] (Ikeya, K. et al. (1989)	
ㅈ 	1382	_
Polypeptide Incyte GenBan SEQ ID NO: Polypeptide ID NO ID	2515666CD1	
Polypeptide SEQ ID NO:	Н	

Analytical Methods and Databases	BLIMPS- PRINTS		BLIMPS-	PRINTS		Ì	BLIMPS-	PRINTS		MOTIFS	BLAST-DOMO		BLAST-	PRODOM	HMMER-PFAM	PROFILESCAN		BLIMPS-	BLOCKS		HMMER	HMMER
Signature Sequences, Domains and Motifs	E-class P450 group I PR00463: S69-L88, T93-S114, S195-S213,	K294-T311, I314-G340, E357-P375, N398-T422, M436-C446, C446-F469	E-class P450 group II PR00464:	G130-A150, K294-V322, T323-G340,	Q352-F372, K392-E407, L408-A423,	S433-C446, C446-F469	P450 superfamily signature PR00385:	A305-V322, T323-D336, A358-F369,	M437-C446, C446-F457	Cytochrome P450: F439-G448	CYTOCHROME P450	DM00022   S43414   18-491: V24-L485	Cytochrome P450 PD000021:	L111-R284, F263-G386, F387-F469	Cytochrome P450: P42-L499	Cytochrome P450 cysteine heme-iron	ligand signature: F415-E468	Cytochrome P450 cysteine heme-iron	ligand signature BL00086:	M436-L467	Transmembrane domain: L14-R32	Signal peptide: M1-C22
Potential Glycosylation Sites			•																			
Potential Phosphorylation Sites	S10 S89 S212 T267 T323 T378	⊣	T486																			
Amino Acid Residues	504																					
Incyte Polypeptide ID	2515666CD1																					
SEQ ID NO:	П																					

Polynucleotide	Incyte	Sequence	İ	Seguence		3′
SEQ ID NO:	Polynucleotide ID	Length	Fragments	Fragments	Position	Position
C1	2515666CB1	1790	1-77,	SZAE00257F1	1356	1790
			891-915	SZAE00059F1	142	770
				2515666H1 (LIVRTUT04)	1	177
				SZAE00073F1	306	1601
				SZAE00189F1	251	871
				SZAE00140F1	843	1582

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LIVRTUT04	251566CB1	2
Library	Project ID	SEQ ID NO:
Representative	Incyte	Polynucleotide

	The second second second second second	
Library	Vector	Library Description
LIVRTUT04	DINCY	Library was constructed using RNA isolated from liver tumor tissue removed from a 50-
		year-old Caucasian male during a partial hepatectomy. Pathology indicated a grade 3-4
		hepatoma, forming a mass. Patient history included benign hypertension and hepatitis.
		Hepatitis B core antigen and hepatitis B surface antigen was present in the patient.

Program	Description	Reference	Parameter Threshold
ABIFACTURA	A program that removes vector sequences and masks ambiguous bases in nucleic acid sequences.	Applied Biosystems, Foster City, CA.	
ABI/PARACEL FDF	A Fast Data Finder useful in comparing and annotating amino acid or nucleic acid sequences.	Applied Biosystems, Foster City, CA; Paracel Inc., Pasadena, CA.	Mismatch <50%
ABI AutoAssembler	A program that assembles nucleic acid sequences.	Applied Biosystems, Foster City, CA.	
BLAST	A Basic Local Alignment Search Tool useful in sequence similarity search for amino acid and nucleic acid sequences. BLAST includes five functions: blastp, blastn, blastx, tblastn, and tblastx.	Altschul, S.F. et al. (1990) J. Mol. Biol. 215:403-410; Altschul, S.F. et al. (1997) Nucleic Acids Res. 25:3389-3402.	ESTs: Probability value= 1.0E-8 or less Full Length sequences: Probability value= 1.0E-10 or less
FASTA	A Pearson and Lipman algorithm that searches for similarity between a query sequence and a group of sequences of the same type. FASTA comprises as least five functions: fasta, tfasta, fastx, tfastx, and ssearch.	Pearson, W.R. and D.J. Lipman (1988) Proc. Natl. Acad Sci. USA 85:2444-2448; Pearson, W.R. (1990) Methods Enzymol. 183:63-98; and Smith, T.F. and M.S. Waterman (1981) Adv. Appl. Math. 2:482-489.	ESTs: fasta E value=1.06E-6 Assembled ESTs. fasta Identity= 95% or greater and Match length=200 bases or greater; fastx E value=1.0E-8 or less Full Length sequences: fastx score=100 or greater
BLIMPS	A BLocks IMProved Searcher that matches a sequence against those in BLOCKS, PRINTS, DOMO, PRODOM, and PFAM databases to search for gene families, sequence homology, and structural fingerprint regions.	Henikoff, S. and J.G. Henikoff (1991) Nucleic Acids Res. 19:6565-6572; Henikoff, J.G. and S. Henikoff (1996) Methods Enzymol. 266:88-105; and Attwood, T.K. et al. (1997) J. Chem. Inf. Comput. Sci. 37:417-424.	Probability value= 1.0E-3 or less
HMMER	An algorithm for searching a query sequence against hidden Markov model (HMM)-based databases of protein family consensus sequences, such as PFAM.	Krogh, A. et al. (1994) J. Mol. Biol. 235:1501-1531; Sonnhammer, E.L.L. et al. (1988) Nucleic Acids Res. 26:320-322; Durbin, R. et al. (1998) Our World View, in a Nutshell, Cambridge Univ. Press, pp. 1-350.	PFAM hits: Probability value=1.0E-3 or less Signal peptide hits: Score=0 or greater

# Table 7 (cont.)

Program	Description	Reference	Parameter Threshold
ProfileScan	An algorithm that searches for structural and sequence motifs in protein sequences that match sequence patterns defined in Prosite.	Gribskov, M. et al. (1988) CABIOS 4:61-66; Gribskov, M. et al. (1989) Methods Enzymol. 183:146-159; Bairoch, A. et al. (1997) Nucleic Acids Res. 25:217-221.	Normalized quality score>GCG-specified "HIGH" value for that particular Prosite motif. Generally, score=1.4-2.1.
Phred	A base-calling algorithm that examines automated sequencer traces with high sensitivity and probability.	Ewing, B. et al. (1998) Genome Res. 8:175-185; Ewing, B. and P. Green (1998) Genome Res. 8:186-194.	
Phrap	A Phils Revised Assembly Program including SWAT and CrossMatch, programs based on efficient implementation of the Smith-Waterman algorithm, useful in searching sequence homology and assembling DNA sequences.	Smith, T.F. and M.S. Waterman (1981) Adv. Appl. Math. 2:482-489; Smith, T.F. and M.S. Waterman (1981) J. Mol. Biol. 147:195-197; and Green, P., University of Washington, Seattle, WA.	Score= 120 or greater; Match length= 56 or greater
Consed	A graphical tool for viewing and editing Phrap assemblies.	Gordon, D. et al. (1998) Genome Res. 8:195-202.	12.
SPScan	A weight matrix analysis program that scans protein sequences for the presence of secretory signal peptides.	Nielson, H. et al. (1997) Protein Engineering 10:1-6; Claverie, J.M. and S. Audic (1997) CABIOS 12:431-439.	Score=3.5 or greater
TMAP	A program that uses weight matrices to delineate transmembrane segments on protein sequences and determine orientation.	Persson, B. and P. Argos (1994) J. Mol. Biol. 237:182-192; Persson, B. and P. Argos (1996) Protein Sci. 5:363-371.	
TMHMMER	A program that uses a hidden Markov model (HMM) to delineate transmembrane segments on protein sequences and determine orientation.	Sonnhammer, E.L. et al. (1998) Proc. Sixth Intl. Conf. on Intelligent Systems for Mol. Biol., Glasgow et al., eds., The Am. Assoc. for Artificial Intelligence Press, Menlo Park, CA, pp. 175-182.	.2.
Motifs	A program that searches amino actd sequences for patterns that matched those defined in Prosite.	Bairoch, A. et al. (1997) Nucleic Acids Res. 25:217-221; Wisconsin Package Program Manual, version 9, page M51-59, Genetics Computer Group, Madison, WI.	:217-221; , page WI.